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- (71) Applicant (for all designated States except US): BLUE MARBLE POLYMERS LIMITED [NZ/NZ]; 61 Wickham Street, Christchurch 8001 (NZ).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): HASTINGS, Maria, Louise [NZ/NZ]; 78B Aorangi Road, Christchurch (NZ). ERRINGTON, John, Gough [NZ/NZ]; 11 Paparoa Street, Christchurch (NZ). HORNSEY, Anya, Jane [NZ/NZ]; 39A Scarborough Road, Christchurch (NZ). CHAPMAN, Timothy, James [NZ/NZ]; 17 Hardwicke Street, Christchurch (NZ). QUINN, Philip, James [GB/NZ]; 259 Innes Road, Christchurch (NZ).

- (74) Agents: WILSON, Kathryn, S. et al.; KPMG Centre, Level 12, 85 Alexandra Street, Private Bag 3140, Hamilton (NZ).
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(54) Title: METHOD FOR THE PRODUCTION OF BIODEGRADABLE FOAMED PRODUCTS













(57) Abstract: A method for the production of a biodegradable foamed product 7 from a base material and a blowing agent 1. The base materials are mixed with a blowing agent and any other required additives 1. The mixture 3, after extrusion 2, is placed in a microwave transparent mould 6 and processed in a microwave via distinct steps. The first step preheats the extrudate to a temperature just below the flash point 6. The second step 7 rapidly heats the extrudate beyond the flash point causing the extrudate to foam in the mould 7. By utilising this method it is possible to produce shaped articles with uniform properties and with packaging properties such as compressibility, resilience and shock absorption.



70 02/14043 A1

# METHOD FOR THE PRODUCTION OF BIODEGRADABLE FOAMED PRODUCTS

#### TECHNICAL FIELD

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The present invention relates to the method for manufacturing biodegradable moulded products. More particularly, the present invention relates to having a matrix based on starch, or materials with similar rheological properties, foamed for use in moulded products. In particular, the method involves two or more distinctive steps during microwave processing, giving a product with improved packaging properties including resilience, compressibility and shock absorption.

#### **BACKGROUND ART**

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The area of starch based biodegradable foamed materials is widely discussed in the prior art. In particular, US patent number 6,168,857 has a detailed discussion that may be referenced in relation to this patent.

#### **Containers and Packaging**

Articles such as sheets, films and packaging moulds made from materials such as paper, paperboard, plastic, polystyrene, and even metals are used in enormous quantity. This can take the form of printed materials, mats, containers, separators, dividers, envelopes, lids, tops, cans, and other packaging materials. Advanced processing and packaging techniques presently allow an enormous variety of liquid and solid goods to be stored, packaged, or shipped while being protected from harmful elements.

Containers and other packaging materials protect goods from environmental influences and distribution damage, particularly from chemical and physical influences. Packaging helps protect an enormous variety of goods from gases, moisture, light, micro-organisms, vermin, physical shock, crushing forces, vibration, leaking, or spilling.

For the purposes of the discussion, many prior art products and processes are seen as not being environmentally friendly. Wherein for the purposes of the present invention, "environmentally friendly" may be characterised as:

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- Being produced from naturally occurring and renewable, raw materials;
- Manufactured in such a way as to cause minimal deterioration to the environment for example via low energy processing and low emission methods;
  - Producing a product that is biodegradable and not harmful to the environment; and
  - Production whereby the whole process is sustainable.

However it is not intended that this definition be seen as limiting.

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#### The Impact of Traditional Materials

Recently there has been a debate as to which of these materials (e.g., paper, paperboard, plastic, polystyrene, glass, or metal) is most damaging to the environment. Paper, paperboard, plastic, polystyrene, glass, and metal materials each have unique environmental issues that do not meet the definition of "environmentally friendly". These issues can relate to the biodegradability of the material itself or the method of production, for example, high energy use, damaging by-products and emissions.

Another problem with paper, paperboard, polystyrene, and plastic is that each of these requires relatively expensive organic starting materials, some of which are non-renewable, such as the use of petroleum in the manufacture of polystyrene and plastic. Although trees used in making paper and paperboard are renewable in the strict sense of the word, their large land requirements and rapid depletion in certain areas of the world undermines this notion. Hence, the use of huge amounts of essentially non-renewable starting materials in making sheets and articles therefrom cannot be sustained and is not wise from a long-term perspective.

#### **Starch Based Foams**

Recent uses of starches and starch derivatives as the binding agent or sole constituent within moulded articles are known. US 5,095,054 is the parent document for this style of product. The parent patent, and the patents citing this patent, recognise the fact that starch can be foamed and moulded by means of forming what is known in the art as "destructurised starch". In the manufacture of destructurised starch, native starch or starch derivatives are mixed with a wide variety of additives such as plasticisers, and heated, solidified and cooled, typically into a mould.

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EP-707034 and WO95/07693 both use conventional thermal conductive heating processes that do not lend themselves to the production of thick-walled mouldings. Non-homogenous heating occurs when the heating process is reliant on heat conduction as it is difficult to heat the core of the material to the same extent as the exterior. This results in non-uniform foam properties, which is undesirable in protective packaging used for cushioning applications.

A further example includes US 6,168,857 in which the process is only usable in thin walled applications. Thin walled articles are of limited use in terms of protective packaging used for cushioning applications. Thick walled articles are needed where shock absorption properties are required. The method of fashioning articles from sheets used in US 6,168,857 does not allow for the forming of thick sheets.

Another patent US 5,730,824 utilises extrusion to produce foam panels. These panels are then laminated together to form thick sheets, which can be wire cut to varying size shapes. There are limitations in this process due to the expensive capital equipment required for manufacturing. As a result of the expensive equipment, the method necessitates shipping 'air' as the product can only be made in central locations. In addition the shapes are either very limited or costly because they have to be cut out of sheets instead of moulded during the foaming process.

Another example, US 5,801,207, relates to taking foamed starch pieces, placing them in a bag or within layers of sheeting and moulding the pre-expanded peanuts into solid foam-in-place moulds. The limitations are that the foamed peanuts used to make the moulds are very bulky and take up a lot of store space, and again increase expense through having to ship air to the point of use instead of sending dense pellets that can be foamed at point of use. The method is also a complicated procedure for the end-user, as they have to fill and seal bags of foamed peanuts and then mould the bag to the product shape.

10 Two further patents, WO 9,851,466 and US 5,639,518, utilise dielectric heating in processing the starch based materials.

In WO 9,851,466, the dielectric heating proceeds in one step and does not take into account the changing dielectric properties of the material as it heats, nor the relationship between the rheological properties (for example elasticity and viscosity) and the rate of heating. This results in the material not being heated as rapidly and intensely, thus lowering the potential foaming and product resilience.

Patent US 5,639,518 again does not utilise different steps during processing to take account of the changing dielectric rheological properties of the material as it heats up. Two stages are outlined relating to changes in the microwave frequency from low frequency and then high frequency but not with any reference to varying material properties and a rate of heating profile. This frequency change results in a significant increase in processing expense due to more specialised equipment being required.

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In addition, the methods described above often produce foams with varying consistency depending on the shape required and often without the combination of uniform physical and mechanical properties. These properties include density, compressibility, resilience and shock absorption. All of these properties limit the product applications.

A further method for moulding starch-based mixtures into articles involves batch-moulding an aqueous starch mixture between heated dies. The starch binder is preferably initially in an unmodified, un-gelatinised state within the mouldable aqueous mixture. The starch/water mixtures are heated between the moulds to a temperature great enough to gelatinise the starch as well as to remove the majority of the water from the mouldable mixture. The resulting moulded articles can be de-moulded, but are initially very brittle until they have been "conditioned" by placing them in a high humidity chamber for extended periods of time in order to reabsorb moisture. While the foregoing batch moulding process may have some utility, it does not allow for continuous manufacturing as is expensive to run.

Based on the above, there is a perceived difficulty in finding improved methods for manufacturing low cost and environmentally friendly products, which have properties similar to paper, paperboard, polystyrene, or plastic, but are biodegradable and resilient.

An object of the present invention is the provision of the method to produce a foamed product with uniform physical and mechanical properties such as density, compressibility, resilience and shock absorption.

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A further object is the provision of a method and product, which overcome some, or all of the above described disadvantages of existing biodegradable foamed products.

Another object is the production of a biodegradable foamed product and method, which provides the public with a useful alternative to existing methods and products.

#### **DISCLOSURE OF INVENTION**

According to one aspect of the present invention there is provided a method of producing a biodegradable foamed product, the method including the steps of;

- 5 (a) selecting one or more base materials,
  - (b) blending the base materials with one or more additives to form a mixture;
  - (c) extruding the mixture in the presence of water,
  - (d) transferring the extrudate to a mould;
  - (e) heating the extrudate using dielectric heating, characterised by the steps of;
- (f) preheating the extrudate to below the flash point of the blowing agent to achieve a uniform temperature distribution throughout the extrudate;
  - (g) rapid heating of the extrudate through the flash point of the blowing agent thereby forming an expanded foamed product.
- It is the understanding of the applicant that the advantages of this method of introducing energy are based on the fact that with dielectric heating it is possible to realise very high energy densities. In addition, the energy acts on the starch material not merely at the surface, but also penetrates into the starch material, allowing rapid internal heating of the material, which results in the flashing off of the blowing agent. Uniform cell structure is achieved through the optimisation of microwave application such that a uniform electric field is generated within the material.

It is understood by the applicant that the bubble growth is a function of the amount of energy delivered to the blowing agent and of the rheological properties of the material.

25 The viscoelasticity of the material must be such that it can allow the build up of water vapour pressure without rupture or severe shrinkage.

In the preferred embodiment, dielectric heating is used for heating the product as it has been found in practice to provide the most even heating and also the required intensity.

According to a further aspect of the present invention there is provided a method of producing a biodegradable foamed product, the method including the steps of;

(a) selecting one or more base materials,

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- (b) blending the base materials with one or more additives to form a mixture;
- (c) extruding the mixture in the presence of water;
- (d) transferring the extrudate to a mould;
- 10 (e) heating the extrudate using dielectric heating, characterised by the steps of;
  - (f) preheating the extrudate to below the flash point of the blowing agent to achieve a uniform temperature distribution throughout the extrudate;
  - (g) rapid heating of the extrudate through the flash point of the blowing agent thereby forming an expanded foamed product; wherein, the force of the water vapour pressure is higher than the viscous forces of the material.

If the viscosity of the extrudate melt is too low, no water vapour pressure will build up as the material will offer no resistance against which the water vapour pressure can build.

- According to a further aspect of the present invention there is provided a method of producing a biodegradable foamed product, the method including the steps of;
  - (a) selecting one or more base materials,
  - (b) blending the base materials with one or more additives to form a mixture;
  - (c) extruding the mixture in the presence of water;
- 25 (d) transferring the extrudate to a mould;
  - (e) heating the extrudate using dielectric heating, characterised by the steps of;

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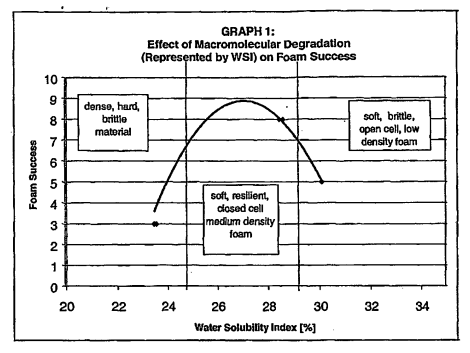
(f) preheating the extrudate to below the flash point of the blowing agent to achieve a uniform temperature distribution throughout the extrudate;

(g) rapid heating of the extrudate through the flash point of the blowing agent thereby forming an expanded foamed product; wherein, the elasticity of the material is such that the blowing agent vapour pressure is high enough to not rupture the extrudate and low enough to prevent severe shrinkage.

If the elasticity is too low during step (g), the water vapour pressure will rupture the extrudate causing a weak open cell structure to be formed. Alternatively, if the elasticity at step (g) is too high, the material will shrink before the structure is able to solidify causing a very high density and hard foam to be formed.

It is the understanding of the applicant that viscosity and elasticity are both functions of a large number of variables including temperature, water content, molecular structure (influenced by the level of molecular degradation), plasticiser content, additive content for example viscosity modifiers and starch gelatinisation.

In tests completed on the material processed using the above method, it has been found that the level of molecular degradation affects the foam success as shown on Graph 1 20 below:



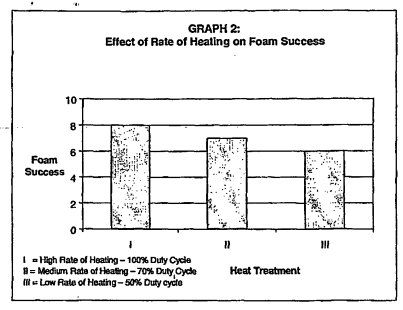
Graph 1 shows that the foam success is influenced by the level of molecular degradation, (as indicated by water solubility index, WSI). Those skilled in the art will appreciate that a correlation therefore exists between viscosity and elasticity of the base material and the level of molecular degradation.

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According to a further aspect of the present invention, the microwave equipment is adjusted such that maximum heating (incorporating power density) is rapidly delivered to the material in stage (g). In a further embodiment, in stage (g) described above, one or more re-tuning steps may be effected depending on the dielectric property changes of the extrudate. Alternatively, separate microwave generators, differently tuned, are used in place of the re-tuning steps.

Varying the intensity of thermal energy as shown in Graph 2 below regulates the foaming success:



In the case of the present invention, it is understood by the applicant that the foam formation involves the nucleation and growth of water vapour bubbles as the result of rapid heating.

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The applicant has found that the more rapid the heating is, the better the foam success. Foam success is measured based on the characteristics of successful packaging foams. These include density, compressibility, resiliency and particle adhesion but it is the combination of these properties that ultimately determines the functionality of the foam. The quantity 'foam success' is used in the application to quantify the overall functionality of the foam.

Hence, given the relationship between viscosity, elasticity and rapid heating, in the preferred embodiment, between step (f) and step (g), the microwave equipment is re-tuned.

In addition to the changes in the viscosity and elasticity, the dielectric properties of the extrudate change considerably between room temperature and around 80-99°C (just below the flash point). It has been found that by re-tuning the microwave equipment, it is rematched to the different dielectric properties of the material at the end of the preheat phase.

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With the equipment/material matching improved, higher energy densities can be achieved and the rate of heating can be optimised in the critical heating region i.e. around 90- 130°C where the foaming occurs i.e. in step (g). More accurate energy density profiles can also be achieved through re-tuning.

For the purposes of the explanation two stages (f) and (g) have been used for an example of the dielectric heating method. This should not be seen as limiting, as further stages are possible before and after reaching the material flash point.

In the preferred embodiment, the microwave radiation is within the standard microwave frequency range 915 MHz to 25 GHz. Preferably, the radiation is in the range easily reached for microwave equipment of 2.0 GHz to 3.0 GHz, more preferably in the standard range for microwave equipment of 2.40 to 2.50 GHz. According to a further aspect of the present invention, the operating frequency of the microwave is held approximately constant during processing. In addition, a power density distribution in the range of 0.01 W/cc to 10.00 W/cc is used. More preferably the energy density is greater than 3.5W/cc.

In a further embodiment, during microwave treatment, the power output of the microwave is varied depending on the desired product characteristics required. Varying the duty cycle of the microwave source is a commonly used means of controlling the effective power output. Alternatively, if more than one microwave generator is used, they may be at different power ratings.

In the above described method, the base material at step (a) is selected from the group consisting of: proteins, starches including cereal, root and tuber starches, modified starches, food residues, biodegradable polymers, and any combination thereof. It can also include materials where the material has rheological properties that are similar to those of starch materials.

In the preferred embodiment, the preferred blowing agent is water. It is understood by the applicant that successful foaming comes from rapid heating, which causes a 'flash-off' of the agent. In the case of water, if the rate of heating is too slow, then the rate of vapour transfer will be higher than rate of heating. This means the water will evaporate, causing no build up in water vapour pressure and hence no water vapour bubbles form or grow to create the 'cells' necessary for a uniform foamed product.

The present invention also provides a method as described above in which the further additives include a blowing agent other than water.

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In a preferred embodiment, a solid blowing agent is added at step (b). Alternatively a liquid blowing agent is added at either step (b) or step (c). The process is not limited to only occur as the result of the change of state of the blowing agent from a liquid into the gaseous phase. There are other mechanisms for foam formation including the thermal degradation of solids to form gases e.g. sublimation or thermally initiated chemical reactions, which produce gases. In a preferred embodiment these gases act as blowing agents.

Other additives can also be included. Typically these additives are selected from a range of biodegradable plasticisers (such as polyvinyl alcohol with various hydrolysis degrees), nucleating agents (such as magnesium silicate, calcium carbonate), processing aids (such as lecithin, mono-glycerides) and any combination thereof.

Further additives with an application dependent function can also be included such as flame retardants, fungus and mould inhibitors, strength adjusting additives, adhesion promoters, viscosity modifiers, fillers and rodent repellents.

According to a further aspect of the present invention, in the above described method, the moisture content, during extrusion, ranges between 10% by weight and 50% by weight.

Preferably the range is from 15% to 30% by weight as this has been found to give the best foaming success in trials. These weight ranges are based upon the total weight of the raw material starting mixture before the microwave heating thereof.

A further option for processing includes a conditioning step following extrusion. The extrudate is conditioned to a pre-determined moisture content via temperature and humidity control. The moisture content is in the range from 5% to 20% by weight as this has been found to give the best foaming success in trials.

During steps (f) and (g), the extrudate is heated dielectrically using moulds to hold the extrudate. The mould container is either completely microwave transparent or at least largely microwave transparent, taking into account the increase in volume of the material, and the fact that the water vapour needs to be expelled from the mould unhindered at the same time.

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A further advantage of the present invention is the low set-up and running cost. The method utilises equipment generally well known and utilises, for example extruders and microwave ovens. Similarly, the materials can be shipped prior to foaming, thus keeping freight costs lower due to the smaller volumes shipped. The customer can then microwave and foam the material thus improving the process efficiency i.e. not paying to ship 'air' in the expanded product and spreading the capital equipment cost.

According to another aspect of the present invention, there is provided a biodegradable foamed product produced by any one of the above described methods.

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According to another aspect of the present invention, there is provided a product produced after step (c).

According to another aspect of the present invention, there is provided a biodegradable product produced after step (f).

#### 5 BRIEF DESCRIPTION OF DRAWINGS

Further aspects of the present invention will become apparent from the following description, which is given by way of example only and with reference to the accompanying drawings in which:

10 Figure 1 is a flow diagram of the process according to a preferred embodiment of the present invention.

#### BEST MODES FOR CARRYING OUT THE INVENTION

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#### Example 1

Referring to the drawing, the method starts with a blend of a starch (as the base material) and other substances mixed together to form a blend (1).

20 An example of a base material blend termed 'A2' for the example, includes:

Material	Proportion
	[% Dry weight]
Tapioca Starch	81.25
Maltodextrin (dextrose equivalent 8-12)	2.5
Pregelatinsed modified waxy maize starch	3.00
Polyvinyl alcohol (degree of hydrolysis = 86-89 mole %)	12.00
Magnesium Silicate	0.25
Lecithin	1.00

However, it will be appreciated that other biodegradable polymers can be used as the base material, provided the rheological properties of the material is similar to those of starch pellets.

The blend (1) is extruded in an extruder (2) of known type at a moisture content of 15% to 30% by weight to achieve an extrudate with the desired rheological properties (3). Such an extruder is, for example, one with a single screw.

The extrudate (3) is cut into pellets or long rods or rolled into final mould ready shapes. The extrudate is conditioned (4) to a moisture content in the range of 12% to 16% by weight at a temperature of 15°C to 40°C. This occurs in 25% to 80% relative humidity in a controlled climate container (4) to minimise case hardening of the extrudate (3). This gives a shelf stable and easily transported product, which can be stored for later use (5).

It will be appreciated by those skilled in the art that other shapes may also be achieved, for example plain or patterned sheets.

The conditioned extrudate (5) is loaded into a microwave-transparent mould in the shape required. For this example, 70g of extrudate (5) is loaded into an 18cm x 8cm x 4cm container (6).

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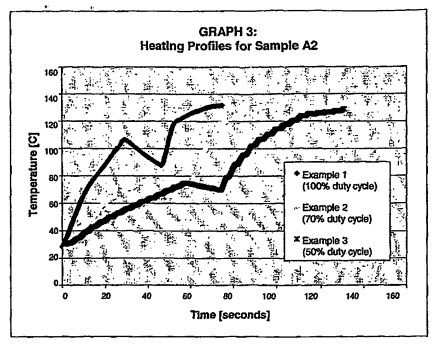
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The mould is placed into a microwave field (6) using 2.45 GHz frequency microwave energy with a variable power output from 100W to 5 kW, for the example set to a 2 kW source. The duty cycle of the microwave is 100%.

For stage A, of the preferred embodiment, the extrudate (5) is pre-heated for less than 30 seconds depending on the tuning of the microwave generator to bring the extrudate (5) to just below the flash point of the water blowing agent. The magnetron (not shown) is adjusted to an energy density of 0.83W/cc whence it is used to heat the extrudate (5) in the mould (6) for less than 30 seconds depending on the tuning of the microwave equipment (not shown). This produces a uniform, resilient, moulded foam block (7).

Re-tuning the magnetron allows for changing dielectric properties in the extrudate as the temperature changes. Better energy densities and more accurate density profiles can thus be achieved. While re-tuning is carried out the frequency remains approximately around 2.45 GHz. A temperature profile is shown for the example in Graph 3 below.



The result of heating at 100% duty cycle is foam success of 8, shown in Graph 2 above. Foam success is measured based on the characteristics of successful packaging foams.

These include density, compressibility, resiliency and particle adhesion but it is the combination of these properties that ultimately determines the functionality of the foam.

#### Example 2

In a second example the same base material, A2 is put through the same steps (1) to (5) described above. During steps (6) and (7) the microwave duty cycle is changed from 100% in example 1, to 70%. Accordingly, the length of time for each step is also varied to account for the altered heating profile. Graph 3 shows the resulting heating profile and Graph 2 shows the resulting foam success is 7.

#### 15 Example 3

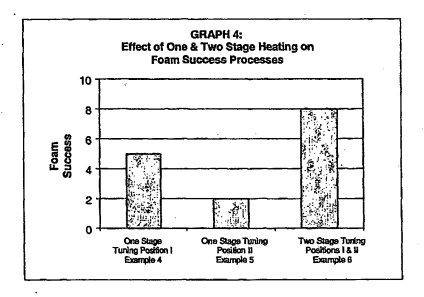
In a third example the same base material, A2 is put through the same steps as example 2 above however a microwave duty cycle 50% is used. The length of time for each step is

again varied to account for the altered heating profile. Graph 3 shows the resulting heating profile and Graph 2 shows the resulting foam success is 6.

#### Example 4

In a fourth example the same base material, A2 is put through the same steps (1) to (5) described in example 1. During steps (6) and (7) the microwave duty cycle is held at 100% and the foaming success is tested with a 60 second single step process with the magnetron tuned to the material at the beginning of the microwave process. The foam success result of 5 is shown in Graph 4 below.

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#### Example 5

In a fifth example the same base material as example 1 is processed as per example 4 above, except that during steps (6) and (7) the foaming success is tested. This is done with a 60 second single step process wherein the magnetron is tuned to the material at the normal tuning used when moving through the flash temperature of the blowing agent. The foam success result of 2 is shown in Graph 4 above.

Example 6

In a sixth example the same base material as example 1 is processed as per example 4 above except that during steps (6) and (7) the foaming success is tested with two 30 second steps wherein the magnetron re-tuned to the material at the end of each step. The foam success result of 8 is shown in Graph 4 above.

Aspects of the present invention have been described by way of example only and it should be appreciated that modifications and additions may be made thereto without departing from the scope thereof as defined in the appended claims.

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#### THE CLAIMS DEFINING THE INVENTION ARE:

1. According to one aspect of the present invention there is provided a method of producing a biodegradable foamed product, the method including the steps of;

- 5 (a) selecting one or more base materials,
  - (b) blending the base materials with one or more additives to form a mixture;
  - (c) extruding the mixture in the presence of water;
  - (d) transferring the extrudate to a mould;
  - (e) heating the extrudate using dielectric heating, characterised by the steps of;
- 10 (f) preheating the extrudate to below the flash point of the blowing agent to achieve a uniform temperature distribution throughout the extrudate;
  - (g) rapid heating of the extrudate through the flash point of the blowing agent thereby forming an expanded foamed product.
- 2. A method of producing a biodegradable foamed product, the method including the steps of;
  - (a) selecting one or more base materials,
  - (b) blending the base materials with one or more additives to form a mixture;
  - (c) extruding the mixture in the presence of water;
- 20 (d) transferring the extrudate to a mould;
  - (e) heating the extrudate using dielectric heating, characterised by the steps of;
    - (f) preheating the extrudate to below the flash point of the blowing agent to achieve a uniform temperature distribution throughout the extrudate;

(g) rapid heating of the extrudate through the flash point of the blowing agent thereby forming an expanded foamed product, wherein the force of the water vapour pressure is higher than the viscous forces of the material.

- 5 3. A method of producing a biodegradable foamed product, the method including the steps of;
  - (a) selecting one or more base materials,
  - (b) blending the base materials with one or more additives to form a mixture;
  - (c) extruding the mixture in the presence of water;
- 10 (d) transferring the extrudate to a mould;
  - (e) heating the extrudate using dielectric heating, characterised by the steps of;
    - (f) preheating the extrudate to below the flash point of the blowing agent to achieve a uniform temperature distribution throughout the extrudate;
    - (g) rapid heating of the extrudate through the flash point of the blowing agent thereby forming an expanded foamed product; wherein, the elasticity of the material is such that the blowing agent vapour pressure is high enough to not rupture the extrudate and low enough to prevent severe shrinkage.
- 4. A method as claimed in any one of claims 1 to 3, wherein the initial water content is in the range from 10% and 50% by weight.
  - 5. A method as claimed in any one of claims 1 to 4, wherein the extrudate is conditioned to a pre-determined moisture content in the range of 5% to 20% by weight prior to step (d).

6. A method as claimed in any one of claims 1 to 5 wherein the mould containers used in microwave treatment are either completely microwave transparent or at least largely microwave transparent.

- 5 7. A method as claimed in any one of claims 1 to 6 wherein the microwave frequency is in the range from 915 MHz to 25 GHz.
  - 8. A method as claimed in any one of claims 1 to 6 wherein the microwave frequency is in the range from 2 GHz to 3 GHz.

- 9. A method as claimed in any one of claims 1 to 6 wherein the microwave frequency is in the range from 2.40 GHz to 2.50 GHz.
- 10. A method as claimed in any one of claims 1 to 6 wherein a single frequency is used for microwave treatment.
  - 11. A method as claimed in any one of claims 1 to 10 wherein the microwave power density is in the range from 0.01 W/cc to 10.00 W/cc.
- 20 12. A method as claimed in any one of claims 1 to 11 wherein the microwave equipment is tuned to take into account variations in the changing dielectric properties of the material.
- 13. A method as claimed in any one of claims 1 to 12 wherein the microwave equipment utilises separate microwave generators, differently tuned, to take into account variations in the changing dielectric properties of the material.

14. A method as claimed in any one of claims 1 to 13 wherein the microwave output is adjusted at any stage during microwave processing.

- 15. A method as claimed in any one of claims 1 to 14 wherein more than one microwave generator is used to give different outputs during microwave processing.
  - 16. A method as claimed in any one of claims 1 to 15 wherein the extrudate is heated in two or more steps during the foaming process.
- 10 17. A method as claimed in any one of claims 1 to 16 wherein the base material at step
  (a) is selected from the group consisting of: proteins, starches including cereal, root
  and tuber starches, modified starches, food residues, biodegradable polymers, and
  any combination thereof.
- 15 18. A method as claimed in any one of claims 1 to 17 wherein the base material at step

  (a) is made up of materials with similar rheological properties to claim 16.
  - 19. A method as claimed in any one of claims 1 to 18 wherein a base material is used at step (d) with rheological properties that match the extruded material.

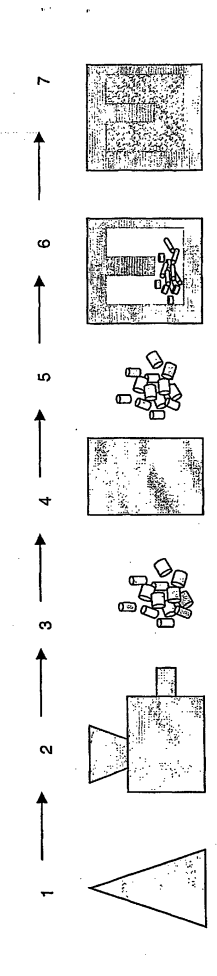
- 20. A method as claimed in any one of claims 1 to 19 wherein the primary blowing agent is water.
- 21. A method as claimed in any one of claims 1 to 20 wherein the primary blowing agent is a solid material that is added at step (b) to form a mixture.

22. A method as claimed in any one of claims 1 to 21 wherein the primary blowing agent is a liquid material that is added at either step (b) or step (c).

- 23. A method as claimed in any one of claims 1 to 22 wherein additives from any of the following are included, being selected from plasticisers, nucleating agents, strength adjusting agents, viscosity modifiers, adhesion promoters, processing aids and fillers.
  - 24. A method as claimed in claim 23 wherein the plasticiser is polyvinyl alcohol.
- 10 25. A method as claimed in claim 24 wherein the nucleating agent is magnesium silicate.
  - 26. A method as claimed in any one of claims 1 to 25 wherein additives are included, said additives included to aid dielectric properties, blowing agents, flame retardant properties, anti-fungal agents, mould inhibitors, and any combination of two or more of these.
  - 27. A biodegradable foamed product produced by the method as claimed in any one of claims 1 to 26.
- 20 28. An intermediate product resulting from steps (c) of the method as claimed in any one of the claims 1 to 26.
  - 29. An intermediate product manufactured after step (f), of the method as claimed in any one of claims 1 to 26.

30. A method of producing a biodegradable foamed product, substantially as hereinbefore described and with reference to the accompanying drawing and examples.

5 31. A biodegradable foamed product, as claimed in any one of claims 27 to 30, substantially as hereinbefore described and with reference to the accompanying drawing and examples.



#### INTERNATIONAL SEARCH REPORT

International application No.

PCT/NZ01/00052

#### CLASSIFICATION OF SUBJECT MATTER Int. Cl. 7: B29C 44/02, B29B 11/10 According to International Patent Classification (IPC) or to both national classification and IPC В. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC: B29C 44/02, 35/08, 35/12. 67/20, 67/22, 69/02, B29B 11/10 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched IPC: AU AS ABOVE Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) DWPI: IPC as above and keywords: Biodegrad+, Starch+, protein+, heat+, microwav+, dielectric+, foam+ expand+, mould+, mold+, extrud+, pellet+ C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category\* Relevant to claim No. EP 692357 A2 (NISSEI KABUSHIKI KAISHA) 17 January 1996 Α 1-31 See whole document WO 98/51466 A (INSTITUT VOOR AGROTECHNOLOGISCH Α 1-31 ONDERZOEK (ATO-DLO) 19 November 1998 See whole document WO 90/08642 A (ADFOAM PTY LIMITED) 9 August 1999 A 1-31 See whole document $|\mathbf{x}|$ See patent family annex Further documents are listed in the continuation of Box C Special categories of cited documents: "T" later document published after the international filing date or "A" Document defining the general state of the art which is priority date and not in conflict with the application but cited to not considered to be of particular relevance understand the principle or theory underlying the invention document of particular relevance; the claimed invention cannot пEп Earlier application or patent but published on or after the international filing date be considered novel or cannot be considered to involve an "L" Document which may throw doubts on priority claim(s) inventive step when the document is taken alone

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1	of the actual completion of the international search y 2001	D	ate of mailing of the international search report  12 Guly 7001	
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#### INTERNATIONAL SEARCH REPORT

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C (Continua	pCT/NZ01/00052 pation). DOCUMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where appropriate, of the relevant passages  Relevant Patent Abstracts of Japan, JP, 11-124459 A (DAEWON PAPTIN FOAM CO LTD)  11 May 1999  See abstract EP 707034 A (WATANABE-ZO SHOTEN CO LTD) 17 April 1996  See whole document				
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### INTERNATIONAL SEARCH REPORT Information on patent family members

International application No. PCT/NZ01/00052

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

	t Document Cited in Search Report			Pate	at Family Member		
EP	692357	AU	24924/95	CA	2154437	JР	8081565
		NZ	272537	US	5639518		
wo	9851466	AU	76767/98				
wo	9008642	AU	48933/90				
JР	11124459	CN	1201729	DE	19736642	FR	2764315
		GB	2326124				
EP	707034	JP	8109278	US	5766749		
							END OF ANNEX

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